we lead up to the 2028 Summer Olympic and Paralympic Games in Los Angeles.

I want to thank my colleagues, Representatives MIKE THOMPSON, DIANA DEGETTE, and BILL JOHNSON for leading this important bipartisan legislation. I would also like to thank Ranking Member RODGERS and all the members and staff of our committee for their efforts to move this legislation forward in a bipartisan manner.

Mr. Speaker, I urge my colleagues to support this legislation, and I reserve the balance of my time.

Mrs. RODGERS of Washington. Mr. Speaker, I yield myself such time as I may consume.

Mr. Speaker, today I rise to speak in support of H.R. 172, the United States Anti-Doping Agency Reauthorization Act, sponsored by Representatives MIKE THOMPSON, BILL JOHNSON, and DIANA DEGETTE.

This bill will reauthorize the U.S. Anti-Doping Agency, USADA, which is the national entity charged with administering anti-doping programs in the United States for Olympic, Paralympic, Pan American, and Parapan American sports.

The U.S. Anti-Doping Agency handles in-competition and out-of-competition testing, results management processes, drug reference resources, and athlete education for all U.S. Olympic and Paralympic Committee-recognized sport national governing bodies, their athletes, and events.

USADA is also the administrator for the Ultimate Fighting Championship Anti-Doping Program.

Reauthorizing this important agency will further the advancement of clean sports, fair games, and positive sportsmanship.

I urge my colleagues to support the bill. We will be hearing from one of my colleagues here in a moment, a colleague on the committee, BILL JOHNSON.

I also want to thank the chairman for working together to get this done today.

Mr. Speaker, I reserve the balance of my time.

Mr. PALLONE. Mr. Speaker, I yield 3 minutes to the gentleman from California (Mr. Thompson) who is the lead sponsor of the legislation.

Mr. THOMPSON of California. Mr. Speaker, I rise today in strong support of H.R. 172, bipartisan legislation reauthorizing the United States Anti-Doping Agency, USADA.

Since 2001, USADA has been recognized by Congress as the official antidoping agency for Olympic, Paralympic, and other sporting competitions in the United States.

The organization conducts drug testing for athletes, manages test results, and pursues bad actors who seek to undermine the principles of clean and fair support through the use of illicit or banned substances.

This important legislation reauthorizes USADA through fiscal year 2030 and provides a slight funding boost to

allow USADA to prepare for the 2028 Olympics in Los Angeles, California.

In addition, this legislation requires USADA to devote a portion of its funding to clean sport initiatives for young athletes and authorizes the Department of Justice and other Federal agencies to cooperate with USADA in the course of its investigations.

I am grateful to my colleagues on the Energy and Commerce Committee for advancing this legislation to the floor, and I urge my colleagues to vote "ves."

Mrs. RODGERS of Washington. Mr. Speaker, I am pleased to yield such time as he may consume to the gentleman from Ohio (Mr. JOHNSON), a colleague and leader on the Energy and Commerce Committee.

Mr. JOHNSON of Ohio. Mr. Speaker, I rise today in support of the U.S. Anti-Doping Agency Reauthorization Act.

I thank my colleague, Representative THOMPSON, for his hard work to get this important legislation across the finish line; and I thank Ranking Member McMorris Rodgers for yielding time.

The United States Anti-Doping Agency, or USADA, has worked hard to ensure that our athletic programs are the best in the world, and also the cleanest. Critical to maintaining that success is ensuring our athletes are competing fairly, without the use of performance-enhancing drugs, which is why I have introduced the U.S. Anti-Doping Agency Reauthorization Act.

USADA must have the resources it needs to ensure the integrity of its programs and advance the American values of sportsmanship and playing by the rules on the global stage.

In addition to funding the agency, this legislation adds a special focus on clean sport training for young athletes and their coaches, and enables USADA to better coordinate with Federal law enforcement.

With the Olympics and other international sporting events just around the corner, I urge my colleagues to join me in supporting this bill.

Mr. PALLONE. Mr. Speaker, I have no additional speakers. I urge support for the legislation, and I yield back the balance of my time.

Mrs. RODGERS of Washington. Mr. Speaker, I have no additional speakers. I urge support, and I yield back the balance of my time.

The SPEAKER pro tempore. The question is on the motion offered by the gentleman from New Jersey (Mr. PALLONE) that the House suspend the rules and pass the bill, H.R. 172.

The question was taken.

The SPEAKER pro tempore. In the opinion of the Chair, two-thirds being in the affirmative, the ayes have it.

Mr. BISHOP of North Carolina. Mr. Speaker, on that I demand the yeas and nays.

The SPEAKER pro tempore. Pursuant to section 3(s) of House Resolution 8, the yeas and nays are ordered.

Pursuant to clause 8 of rule XX, further proceedings on this motion are postponed.

FOOD ALLERGY SAFETY, TREAT-MENT, EDUCATION, AND RE-SEARCH ACT OF 2021

Mr. PALLONE. Mr. Speaker, I move to suspend the rules and pass the bill (S. 578) to improve the health and safety of Americans living with food allergies and related disorders, including potentially life-threatening anaphylaxis, food protein-induced enterocolitis syndrome, and eosinophilic gastrointestinal diseases, and for other purposes.

The Clerk read the title of the bill. The text of the bill is as follows:

S 578

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE.

This Act may be cited as the "Food Allergy Safety, Treatment, Education, and Research Act of 2021" or the "FASTER Act of 2021"

SEC. 2. FOOD ALLERGY SAFETY.

- (a) IN GENERAL.—Section 201(qq)(1) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321(qq)(1)) is amended by striking "and soybeans" and inserting "soybeans, and sesame".
- (b) EFFECTIVE DATE.—The amendment made by subsection (a) shall apply to any food that is introduced or delivered for introduction into interstate commerce on or after January 1, 2023.

SEC. 3. REPORT TO CONGRESS.

- (a) REPORT.—Not later than 18 months after the date of enactment of this Act, the Secretary of Health and Human Services (referred to in this section as the "Secretary") shall submit to the Committee on Health, Education, Labor, and Pensions of the Senate and the Committee on Energy and Commerce of the House of Representatives a report that includes—
- (1) descriptions of ongoing Federal activities related to— $\,$
- (A) the surveillance and collection of data on the prevalence of food allergies and severity of allergic reactions for specific food or food ingredients, including the identification of any gaps in such activities;
- (B) the development of effective food allergy diagnostics;
- (C) the prevention of the onset of food allergies;
- (D) the reduction of risks related to living with food allergies; and
- (E) the development of new therapeutics to prevent, treat, cure, and manage food allergies; and
- (2) specific recommendations and strategies to expand, enhance, or improve activities described in paragraph (1), including—
- (A) strategies to improve the accuracy of food allergy prevalence data by expanding and intensifying current collection methods, including support for research that includes the identification of biomarkers and tests to validate survey data and the investigation of the use of identified biomarkers and tests in national surveys;
- (B) strategies to overcome gaps in surveillance and data collection activities related to food allergies and specific food allergens; and
- (C) recommendations for the development and implementation of a regulatory process and framework that would allow for the timely, transparent, and evidence-based modification of the definition of 'major food allergen' included in section 201(qq) of the Federal Food, Drug and Cosmetic Act (21 U.S.C. 321(qq)), including with respect to—

(i) the scientific criteria for defining a food or food ingredient as a "major food allergen" pursuant to such process, including recommendations pertaining to evidence of the prevalence and severity of allergic reactions to a food or food ingredient that would be required in order to establish that such food or food ingredient is an allergen of public health concern appropriate for such process; and

(ii) opportunities for stakeholder engagement and comment, as appropriate, in considering any such modification to such definition.

(b) PUBLICATION.—The Secretary shall make the report under subsection (a) available on the internet website of the Department of Health and Human Services.

The SPEAKER pro tempore. Pursuant to the rule, the gentleman from New Jersey (Mr. PALLONE) and the gentlewoman from Washington (Mrs. RODGERS) each will control 20 minutes.

The Chair recognizes the gentleman from New Jersey.

GENERAL LEAVE

Mr. PALLONE. Mr. Speaker, I ask unanimous consent that all Members may have 5 legislative days in which to revise and extend their remarks and include extraneous material on S. 578.

The SPEAKER pro tempore. Is there objection to the request of the gentleman from New Jersey?

There was no objection.

Mr. PALLONE. Mr. Speaker, I yield myself such time as I may consume.

Mr. Speaker, I rise today in support of the Food Allergy Safety, Treatment, Education, and Research Act, also known as the FASTER Act.

An estimated 32 million Americans, including approximately one in every 13 children, are affected by food allergies. These allergies can pose significant risks, particularly when inaccurate food labels fail to warn consumers about the presence of some of these allergens.

Under current law, eight allergens are considered major food allergens. These allergens include milk, eggs, shellfish, tree nuts, wheat, peanuts, and soybeans. Due to their status as major food allergens, manufacturers must clearly state the presence of any of these ingredients on labels.

Notably missing from this list of allergens is sesame. Sesame is considered an allergen of growing concern. While its prevalence has more than doubled over the last decade, it is not required to be listed as an allergen on food packaging. In fact, in some cases, a food may contain sesame, but the ingredient won't be listed at all on the labels, instead being referred to through nonspecific terms such as "spices" or words that may not be easily recognized by consumers as containing sesame, such as tahini.

While many may not recognize the significance of a simple ingredients label, for many families, a lack of clarity on ingredients could mean life or death for those who are allergic to sesame. Clearly, this information should be prominently featured on all packaged food labels.

This is an issue we have been working on for quite some time. I previously

introduced a bill several years ago that would list sesame as a major food allergen, and although the Food and Drug Administration opened a docket to solicit feedback about sesame labeling, the agency has not been able to require the listing of sesame due to overly long regulatory processes.

So today, Mr. Speaker, we are bypassing these regulatory delays and taking action. The appropriatelynamed FASTER Act would quickly move this process along by recognizing sesame as a major food allergen and requiring its listing on new food labels after a phase-in process.

The bill will also require FDA to report recommendations on how we can make additional improvements to protect individuals with food allergies, including ways to add additional major food allergens.

So I want to thank Representative MATSUI for her tireless work on this bill. She is the prime sponsor. We came so close to getting this over the finish line in the last Congress, and I am glad that today we are sending this bill to President Biden for his signature. I am proud to support the legislation. I encourage all Members to vote for it.

Mr. Speaker, I reserve the balance of my time.

Mrs. RODGERS of Washington. Mr. Speaker, I yield myself such time as I may consume.

Mr. Speaker, I rise today in support of S. 578, the Food Allergy Safety, Treatment, Education, and Research Act. This legislation is bipartisan, a companion of H.R. 1202, that was led by Representatives MCHENRY, GONZALEZ, and MATSUL.

S. 578 will designate sesame as a major food allergen. This means that, with the enactment of this legislation, manufacturers would have to list this ingredient on the food packaging label of products containing sesame.

Recent studies indicate that sesame allergies are of growing concern in the United States, with a prevalence rate on par with allergies to soy and fish, which are both listed as major allergens under the Federal Food, Drug, and Cosmetic Act.

This commonsense legislation will provide consumers with the information they need to protect themselves and their families from certain dangerous and life-threatening allergic reactions.

Mr. Speaker, I urge my colleagues to support this legislation, and I reserve the balance of my time.

Mr. PALLONE. Mr. Speaker, I yield such time as she may consume to the gentlewoman from California (Ms. MATSUI) who is the House sponsor of the legislation.

Ms. MATSUI. Mr. Speaker, I rise today in support of two of my bills being considered under suspension today, the FASTER Act and the TRANSPLANT Act.

The Food Allergy Safety, Treatment, Education, and Research Act, FAST-ER—which wasn't as fast as I wanted it to be, but this act will help to improve the safety of more than 32 million Americans, including 5.6 million children, living with potentially lifethreatening food allergies.

Under current law, mandatory labeling is required for major food allergens recognized by the FDA like milk, eggs, and peanuts.

My grandson, Robby, has a peanut allergy, and for families like mine, accurate food ingredient labels are vital to making safe and healthy choices. The time we have spent reading the labels and having discussions about whether he can go to a birthday party or not, or go to camp or not, and having friends over, it was just heartbreaking. We need to have those labels be clear.

Critically, the FASTER Act extends these labeling protections to nearly 1.6 million Americans allergic to sesame by requiring sesame to be included as an ingredient on a packaged food label.

The bill also expands the research necessary to find new treatments and is an important step in the right direction to finding an eventual cure for food allergies.

Today is a testament to the hard work of thousands of grassroots advocates who sent emails, made calls, and visited Members of Congress and staff to build support and make sesame the ninth allergen to be labeled under law.

The outpouring of support was incredible. The FASTER Act will truly make a difference for those living with potentially life-threatening food allergies, and we are proud that it can now be sent to President Biden's desk.

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I also rise today in support of H.R. 941, the TRANSPLANT Act, my legislation to reauthorize the C.W. Bill Young Cell Transplantation Program and the National Cord Blood Inventory for another 5 years.

Every 3 minutes, someone is diagnosed with blood cancer. For patients and families facing these fatal diseases, a bone marrow or cord blood transplant may be the best treatment or only potential for a cure.

Congress has long recognized the need to coordinate lifesaving transplants between patients and unrelated donors at the national level and has shown strong bipartisan support over the years for the program.

We must continue to encourage donors and give these patients with otherwise fatal blood cancers a second chance at life. That is why I joined with Representative BILIRAKIS to introduce the TRANSPLANT Act.

This past year, there has been a new sense of urgency for this timely reauthorization. We have seen how Be The Match's status, as the designated operator of the national program, has helped bone marrow couriers continue to facilitate transplants during the pandemic.

We must act swiftly to preserve this critical designation and ensure patients with otherwise fatal blood cancers continue to have access to transplants, both during and after the current public health crisis.

I urge my colleagues to support this legislation today so we may further prevent any lapse in funding. I support both bills.

Mrs. RODGERS of Washington. Mr. Speaker, I yield 3 minutes to the gentleman from North Carolina (Mr. McHenry), one of the leaders of this legislation and the leader of the Financial Services Committee.

Mr. McHENRY. Mr. Speaker, I thank Mrs. Rodgers, my classmate and a member of the Energy and Commerce Committee for yielding. It is an amazing thing to be with you here today, and it is an amazing thing to be here today to talk about this important bill.

As the lead Republican cosponsor of the House companion to S. 578, I rise in support of the FASTER Act.

Millions of Americans suffer from life-threatening food allergies. More than 1.5 million Americans are allergic to sesame, in particular, yet there is no current requirement to include the ingredient on product labels. This legislation would declare it the ninth major allergen to be recognized by the U.S. Food and Drug Administration and update laws to require the labeling of sesame.

This bill would also require the Secretary of Health and Human Services to regularly review promising food allergy treatments and research. This is a major bipartisan priority. These efforts will help slow this rapidly growing disease and ultimately find and fund a cure.

Finally, the FASTER Act establishes a scientific process and framework for establishing additional allergens covered by the Federal Food, Drug, and Cosmetic Act.

I am proud to serve as cofounder and co-chair of the newly formed Congressional Food Allergy Research Caucus, along with Congresswoman DORIS MATSUI. We recognize there is more we can do to help those 32 million Americans, including many who are children who suffer from food allergies.

We can and we should do more to increase funding into research, therapies, and treatments for food allergies. Sending this legislation to the President's desk would be a major first step to achieving our goal of improving treatment opportunities.

I urge my colleagues on both sides of the aisle to vote "yes" on this bill.

Mr. PALLONE. Mr. Speaker, I urge my colleagues to support this bill, S. 578, the FASTER Act, and I yield back the balance of my time.

Mrs. RODGERS of Washington. Mr. Speaker, I have no additional speakers, and I yield back the balance of my time.

The SPEAKER pro tempore. The question is on the motion offered by the gentleman from New Jersey (Mr.

PALLONE) that the House suspend the rules and pass the bill, S. 578.

The question was taken.

The SPEAKER pro tempore. In the opinion of the Chair, two-thirds being in the affirmative, the ayes have it.

Mr. BISHOP of North Carolina. Mr. Speaker, on that I demand the yeas and navs.

The SPEAKER pro tempore. Pursuant to section 3(s) of House Resolution 8, the yeas and nays are ordered.

Pursuant to clause 8 of rule XX, further proceedings on this motion are postponed.

JOHN LEWIS NIMHD RESEARCH ENDOWMENT REVITALIZATION ACT OF 2021

Mr. PALLONE. Mr. Speaker, I move to suspend the rules and pass the bill (H.R. 189) to amend the Public Health Service Act to provide that the authority of the Director of the National Institute on Minority Health and Health Disparities to make certain research endowments applies with respect to both current and former centers of excellence, and for other purposes.

The Clerk read the title of the bill. The text of the bill is as follows:

H.R. 189

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE.

This Act may be cited as the "John Lewis NIMHD Research Endowment Revitalization Act of 2021".

SEC. 2. RESEARCH ENDOWMENTS AT BOTH CURRENT AND FORMER CENTERS OF EXCELLENCE.

Paragraph (1) (beginning with "(1) IN GENERAL") of section 464z–3(h) of the Public Health Service Act (42 U.S.C. 285t(h)) is amended to read as follows:

"(1) IN GENERAL.—The Director of the Institute may carry out a program to facilitate minority health disparities research and other health disparities research by providing for research endowments—

"(A) at current or former centers of excellence under section 736; and

"(B) at current or former centers of excellence under section 464z-4.".

The SPEAKER pro tempore. Pursuant to the rule, the gentleman from New Jersey (Mr. PALLONE) and the gentlewoman from Washington (Mrs. RODGERS) each will control 20 minutes.

The Chair recognizes the gentleman from New Jersey.

GENERAL LEAVE

Mr. PALLONE. Mr. Speaker, I ask unanimous consent that all Members may have 5 legislative days in which to revise and extend their remarks and include extraneous material on H.R. 189.

The SPEAKER pro tempore. Is there objection to the request of the gentleman from New Jersey?

There was no objection.

Mr. PALLONE. Mr. Speaker, I yield myself such time as I may consume.

Mr. Speaker, I rise today in support of H.R. 189, the John Lewis National Institute on Minority Health and Health Disparities Research Endowment Revitalization Act of 2021. This legislation would resume grants to minority academic institutions that fell out of eligibility for funding through the NIMHD Research Endowment Program.

By allowing those schools to resume eligibility, we will once again help these institutions conduct critical research into minority health disparities.

COVID-19, Mr. Speaker, has further exacerbated many of the inequities that minority communities experience when interacting with the healthcare system, inequities that we know existed long before the pandemic. In order to address the inequities in our healthcare system and in our society, we must confront them head-on and work together to eliminate them.

By supporting NIMHD and the academic institutions funded through it, we are helping to advance minority health disparity research and strengthen the diversity of the scientific workforce by recruiting and retaining individuals underrepresented in these fields.

This bill is a step toward progress and an equitable public health system. This bill is named after our former colleague, the late and great Congressman John Lewis from Georgia, who introduced this legislation last Congress. He was a dear friend and a longtime champion of eliminating disparities across the board, and he is certainly missed.

I want to thank my colleagues, the two sponsors, Representatives Barragan and Carter, for leading the effort on this legislation this year. This is truly bipartisan.

Mr. Speaker, I urge my colleagues to support this bill, and I reserve the balance of my time.

Mrs. RODGERS of Washington. Mr. Speaker, I yield myself such time as I may consume.

Mr. Speaker, I rise today in support of H.R. 189, the John Lewis NIMHD Research Endowment Revitalization Act, which was introduced by my colleagues, Representatives Barragan, Carter, and Taylor.

This bill will authorize the National Institute on Minority Health and Health Disparities to award research grants to current and former centers of excellence that conduct research on minority health disparities.

Health inequities are disproportionately experienced by minority populations, and these disparities can have adverse impacts on health outcomes, economic opportunities, and overall quality of life. The current COVID-19 pandemic has only underscored these disparities, which is why this bill is so important.

Continued support of these centers of excellence is critical in advancing minority health, addressing health inequities, and expanding educational and training opportunities for those interested in further advancing research in this space.

I would like to thank my colleagues and especially my colleague on the Energy and Commerce Committee on the